

potential, stomatal regulation

 Abbreviations: DOY: day of the year; DAS: day after sowing; TUE: transpiration use efficiency; SF: sap flow; LAI: green leaf area index; PAR: photosynthetically active radiation; VPD: vapor pressure deficit; An: net leaf photosynthesis; E: leaf transpiration; ψleaf: leaf water potential; ψsunlitleaf: leaf water potential of sunlit 48 leaf; ψ_{shadedleaf}: leaf water potential of shaded leaf; K_{soil}: hydraulic conductance of soil; K_{root}: root hydraulic conductance; Kstem: stem hydraulic conductance; ψsoil_effec: effective soil water potential; ψdifference: 50 difference between effective soil water potential and sunlit leaf water potential; Ksoil root: root system 51 hydraulic conductance (includes soil and root hydraulic conductance); Ksoil_plant: whole plant hydraulic conductance (includes below and aboveground components).

1.Introduction

 Maize (*Zea mays L.*) is a major staple crop throughout the world. Drought stress, which negatively affects crop growth and yield, is of increasing concern in several important maize cultivating regions (Daryanto et al., 2016). Increases in frequency and severity of drought events due to climate change have been recently reported (IPCC, 2022). Thus, field observations and understanding on how maize responds to water stress are necessary to suggest promising traits for breeding programs (Vadez et al., 2021) as well as irrigation schemes (Fang and Su, 2019; Q. Cai et al., 2017). Improved understanding of crops' response to drought can be incorporated into soil-crop models (e.g. crop modelling and soil-vegetation-atmosphere transfer modelling).

 Stomatal regulation is often considered as a key aboveground hydraulic variable in regulating water use 63 of crops. Maize was-is considered as isohydric plant in which stomata are closed in response to sensing 64 drought conditions to maintain leaf water potential (ψ_{leaf}) above critical levels ($\psi_{\text{threshold}}$ or minimum ψ_{leaf}) (Tardieu and Simonneau, 1998). Investigations of how stomatal controls differ among species and genotypes commonly observed minimum ψleaf or analyzed of genetic variability of stomatal control in response to varying soil water content. Analyzing measurements of ψleaf from 400 lines of maize of tropical and European origins under greenhouse and growth chamber conditions, Welcker et al. (2011) reported 69 values of minimum ψ_{leaf} from -0.8 to -1.5 MPa, indicating genetic variability of stomatal responses. The isohydric behavior is due to different mechanisms including hydraulic and/or chemical (e.g. abscisic acid

 [ABA]) signals (Tardieu, 2016). The degree to which these underlying mechanisms interact and differ among genotypes and/or environmental scenarios in explaining the stomatal regulation is still debated 73 (Tardieu, 2016;_r Hochberg et al., 2018). Field evidence in variation of the minimum ψ_{leaf} of maize due to 74 soil water availability and soil hydraulics is rarely reported.

 Water flow along the soil-plant-atmosphere continuum is determined by a series of hydraulic 76 conductivities and gradients in water potential. Hydraulic conductance of soil (Ksoil), root hydraulic 77 conductance (K_{root}), and stem hydraulic conductance (K_{stem}) determine water potential from soil to root 78 and root xylem water, and thus magnitude of ψ_{leaf}. There are two main resistances to water flow from the 79 soil to the shoot, namely the soil and the root resistances, often expressed as their inverse, Ksoil and Kroot (Nguyen et al., 2020; Cai et al., 2018). In wet soils, the soil hydraulic conductivity is much higher than that of roots, and water flow is mainly controlled by root hydraulic conductivity (Hopmans and Bristow, 2002; Draye et al., 2010). It is well-known that a decrease in soil matric potential and soil hydraulic conductivity triggers stomatal closure and thus results in reduction in transpiration rate (Sinclair and Ludlow, 1986; Carminati and Javaux 2020; Abdalla et al., 2021). For the root water uptake and controlling stomata, the location where soil and roots are in close contact (rhizosphere) is most important, because when this thin layer of rhizosphere is disconnected (i.e. soil-root contact is lost), the water movement from soil toward the roots is reduced, which might trigger stomatal closure to maintain hydraulic integrity of plant (Carminati et al., 2016; Rodriguez-Dominguez and Brodribb, 2019; Abdalla et al., 2022). The magnitude of the drop of water potential between bulk soil and soil-root interface increases considerably at different levels of soil dryness for different soil types (Carminati and Javaux, 2020; Abdalla et al., 2022). Hydraulic limits in the soil (Carminati and Javaux, 2020), or in the root–soil interface [as measured for olive trees by Rodriguez-Dominguez and Brodribb, 2019 or tomato (Abdalla et al., 2022)], or in the root properties (Bourbia et al., 2021; Cai et al., 2022; Nguyen et al., 2020; Cai et al., 2018) or due to both soil textures and root phenotypes (Cai et al., 2022b) emphasized the importance of belowground hydraulics (Carminati and

 Javaux, 2020). However, also the shoot hydraulic conductance could be limiting in some crop plants (Gallardo et al., 1996) or in trees (Domec and Pruyn, 2008; Tsuda and Tyree, 1997). Stomatal conductance 97 and shoot hydraulic conductance showed close links to each other in pine trees (Hubbard et al., 2001). This summary illustrates three points: (i) current studies have often focused either on above or on below hydraulic limits, but rarely consider both (ii) it is unclear the roles and relations of soil hydraulic properties to root and plant hydraulic conductance (thus influences on stomatal conductance) (iii) the role of different hydraulic processes across the soil - plant - atmosphere continuum i.e. soil to roots, stem, and soil-plant hydraulic conductance in controlling stomatal conductance remains unclear.

 Simultaneous measurements of atmospheric conditions (light intensity and vapor pressure deficit), leaf water potential, and transpiration rates, coupled with measurements of root, stem and whole soil-plant hydraulic conductance, root architecture, and soil water potential distribution could reveal the relative importance of rhizosphere, shoot and root growth, and hydraulic conductance vulnerability, especially under progressive soil drying at field conditions (Carminati and Javaux, 2020; Tardieu et al., 2017). For the 108 soil water conditions, soil texture and hydraulic characteristics are very important that because they influence soil water movement and thus affect infiltration, surface and sub-surface runoff, and ultimately plant available soil water (Vereecken et al., 2016). Soil texture properties, characterized by different fractions of clay, silt, and sand particles, are important drivers in determining the soil water retention properties (Scharwies and Dinneny, 2019; Stadler et al., 2015; Zhuang et al., 2001). Soil with higher water holding capacity (here the silty soil with low stone content) have a larger amount of plant available water which in turn enables crops to better meet the evaporative demand and facilitates better crop growth as compared to the soil with high stone content (Nguyen et al., 2020; Cai et al., 2018). Estimations of hydraulic conductance (different organs and whole plant hydraulic conductance) were done for crop plants and maize mainly under controlled environment or pot conditions e.g. for different species and genotypes during soil drying (Sunita et al., 2014; Choudhary and Sinclair, 2014; Abdalla et al., 2022; Meunier et al.,

 2018; Wang et al., 2017; Li et al., 2016) or various species and genotypes together with different soil textures (Cai et al., 2022a), or soil texture with different vapor pressure deficit (VPD) (Cai et al., 2022b). Compared to the substantial effect of soil texture, there was no evidence of an effect of VPD on both soil– plant hydraulic conductance and on the relation between canopy stomatal conductance and soil–plant hydraulic conductance in pot-grown maize (Cai et al., 2022b). Contrast results were found in winter wheat where plant hydraulic conductance increased with rising VPD for some genotypes in wet conditions (Ranawana et al., 2021). Vadez et al., (2021) examined the effects of soil types together with increasing VPD on transpiration efficiency (TE) and yield under pot conditions for several C⁴ species (maize, sorghum, and millet). The interpretation of differences in TE was attributed to soil types, more specifically, to the differences in soil hydraulic properties and soil hydraulic conductance. However, experimental evidence linking root hydraulics to stomatal regulation was lacking in these two Vadez's studies (Vadez et al., 2021). Extrapolation and use of results obtained in pots or under greenhouse conditions to the field scale are difficult due to the fact that soil substrates in pots might not represent natural soil in the field (Passioura, 2006). There is often greater evaporative demand and considerable fluctuation and interactions of climatic variables in the field as compared to experiments under controlled or semi-controlled conditions. Recent field studies have aimed at quantification of root hydraulic conductance and it's linkages with crop growth (leaf area and biomass) under different soil types (in wheat Cai et al., 2017; Cai et al., 2018; Nguyen et al., 2020 or maize in Nguyen et al., 2022; Jorda et al., 2022). However, field studies that consider both below (soil-root hydraulic conductance) and above (stem hydraulic conductance), or soil-plant hydraulic conductance (includinges below and above-ground parts) and their roles in stomatal regulation as well as crop growth (leaf area and biomass) are rarely carried out.

 This study aims at further understanding of the hydraulic linkages between soil and plant and responses of plants to drought stress in relation to root: shoot growth characteristics at field scale. We hypothesize that, in field-grown maize, (1) soil-plant hydraulic conductance depends on soil hydraulic properties,

2. Materials and methods

2.1. Location and experimental set-up

 We carried out a field experiment at two rhizotron facilities in Selhausen, North Rhine-Westphalia, Germany (50°52'N, 6°27'E). The field is slightly inclined with a maximum slope of around 4°. One rhizotrone facility was located upslope (F1) with around 60% gravel by weight in the 10-cm topsoil while the second rhizotrone facility was at downslope (F2) with silty soil (stone content is around 4% by weight).

 Each experimental siterhizotrone facility was divided into three subplots of 7.25 m by 3.25 m: two rainfed plots (P1, P2), and one irrigated plot (P3). In rainfed plots P1, other sowing densities and dates were used than in the other plots and we excluded therefore these plots. Silage maize *cv.* Zoey was sown on 4 May 162 and 8 May in 2017 and 2018, respectively, with a plant density of 10.66 seeds m^2 (Figure 1a; Table 1). Detailed information of crop management practices is provided in Table 1.

[Insert Table 1 here]

2.2. Water applications

 Weather variables (global radiation, temperature, relative humidity, precipitation, and wind speed) were recorded every 10 minutes by a nearby weather station (approx. 100 m from the experiment). Drip lines (T-Tape 520-20-500, Wurzelwasser GbR, Müzenberg, Germany) were installed for irrigation at 0.3 m intervals parallel to the crop rows. In 2017, maize received a total amount of 230 mm precipitation during the growing period (136 days). Average, minimum and maximum daily air temperature were 17.6, 8.3, and 25.3 °C, respectively (Fig. 1b). The crop on P3 was irrigated (in total 130 mm) every 5-7 days (in total 10 times) using 13 mm of irrigation water per event between mid June to end of August for the irrigated plots (2017F1P3 and 2017F2P3) (Fig. 1b). In 2018, average, minimum, and maximum daily air temperature were 19.2, 10.85, and 27.3 °C, respectively (Fig. 1b) and exceeded those of 2017. Characterized by exceptionally hot and dry weather conditions, the summer season 2018 can be classified as an extreme year with respect 176 to plant growth at our experimental locationsite. Maize experienced high temperatures and VPD, especially around tasseling and silking. In 2018, only 91.3 mm of rain were recorded in the growing period of 2018 (107 days). The maize crop was irrigated every 5-7 days (in total 13 times), with a total amount of irrigation of 257 mm and 239 mm between mid- June and mid- August for the irrigated plots 2018F1P3 and 2018F1P32018F2P3, respectively (Fig. 1d). In contrast to 2017, the rainfed plot in the stony soil 181 (2018F1P2) had to be irrigated (in total 66 mm) \overline{m} four times (using 13, 22, 13, and 18 mm, respectively) 182 to avoid a crop failure due to severe drought (Fig. 1d). Detailed estimates of irrigation amount and 183 intervals could be found in Nguyen et al., (2022a).

[Insert Figure 1 here]

2.3. Measurements

2.3.1. Soil water measurement and root growth

 At soil depths of 10, 20, 40, 60, 80, and 120 cm, MPS-2 matrix water potential and temperature sensors (Decagon Devices Inc., UMS GmbH München, Germany) were installed to measure half-hourly soil water 189 potential and soil temperature. The range of the water potential measurements is form-from -9 kPa to approximately -100000 kPa (pF 1.96 to pF 6.01). In addition to MPS-2, soil water potential was measured by pressure transducer tensiometers (T4e, UMS GmbH, München, Germany) where the minimum detectable suction is -85 kPa to +100 kPa. A detailed description of sensor installation, calibration and data post processing can be found in Cai et al., (2016).

 Minirhizotubes (7 m long clear acrylic glass tubes with outer and inner diameters of 6.4 and 5.6 cm, respectively) were installed horizontally at six different depths of 10, 20, 40, 60, 80, and 120 cm below the soil surface in each facility. There are three replicate tubes at each depth, accounting for 54 tubes in each facility. Root measurements were taken manually by Bartz camera (Bartz Technology Corporation) (23 June 2017 – 12 September2017) and VSI camera (Vienna Scientific Instruments GmbH) (08 June 2017 – 22 199 June 2017) in 2017 while only VSI was used in 2018 (23 May2018 - 23 August 2018). Root images were taken at 20 fixed positions from the left- and right-hand sides of each tube weekly (or biweekly) during the 201 growing seasonsRoot images were repeatedly taken from both left and right sides at 20 locations along 202 horizontally installed minirhizotubes. The root images were analyzed by automated minirhizotube image analysis pipeline for segmentation and automated feature extraction (Bauer et al., 2021). Two-dimensional 204 root length density (RLD, in units of cm cm⁻²) was estimated from the total root length observed in the image and the image surface area. The overview of camera system, minirhizotube images acquisition, and 206 post-processing of the root data were described in detail in Bauer et al. (2021) and Lärm et al., (2023).

 2.3.2. Crop growth, **leaf gas exchange, leaf water potential, and sap flow measurements measurement**

221 **2.3.3. Leaf gas exchange, leaf water potential, and sap flow measurements**

222 Hourly leaf stomatal conductance (Gs), net photosynthesis (An), and leaf transpiration (E) were measured 223 every two weeks under clear sky conditions. Observations from 8 AM to 5 PM on four days and from 10 224 AM to 4 PM on six days were carried out in 2017. In 2018, measurements were carried out on 6 days from 225 8 AM to 7 PM and on 5 days from 10 AM to 4 PM (Nguyen et al., 2022a). The Gs, An, and E of two sunlit 226 leaves (uppermost fully developed leaves) and one shaded leaf of different plants were measured at 227 steady-state using a LICOR 6400 XT device (Licor Biosciences, Lincoln, Nebraska, USA). After leaf gas 228 exchange measurements, leaves were quickly detached using a sharp knife to measure ILeaf water 229 potential (ψ_{leaf}) was measured with a digital-pressure chamber (SKPM 140/ (40-50-80), Skye Instrument 230 Ltd, UK) with the working air pressure ranging from 0 to 35 bars. To study the diurnal course of ψ_{leaf} under 231 dry and re-wetted soil conditions, in 2018, measurements were undertaken for three additional days with

2.4. Calculation of total root length, root system conductance, stem, and whole plant hydraulic conductance

 To estimate the total root length from minirhizotubes, we adopted the option 2 which was described in 247 Cai et al., (2017). Total root length per square meter soil surface area within each soil layer (m m⁻²) was computed by multiplying the root length density with the corresponding soil layer thickness. The root length density was determined in each depth by dividing the measured root length per minirhizotron 250 image by the assumed volume the roots would have occupied in absence of the tube, i.e., W $*$ L $*$ tube radius (see Cai et al., 2017).

 Following Nguyen et al., (2020), the effective soil water potential was calculated based on hourly measured soil water potential (ψi) and normalized root length density at six depths (10, 20, 40, 60, 80, and 120 cm) (NRLDi), and soil layer thickness (Δzi) in the soil profile (Equation 1).

$$
\psi_{soil_effec} = \sum_{i=1}^{N} \psi_i NRLD_i \,\Delta z_i \tag{1}
$$

255 We followed Ohm's law analogy by dividing the hourly sap flow by the difference between effective soil 256 water potential and shaded leaf water potential to estimate root system conductance (Ksoil_root - Equation 257 2), between shaded leaf water potential and sunlit leaf water potential to estimate stem hydraulic 258 conductance (K_{stem} - Equation 3), and between effective soil water potential and sunlit leaf water potential 259 to estimate whole plant hydraulic conductance (Ksoil_plant - Equation 4).

$$
K_{\text{solid} \text{root}} = \text{Sapflow} / (\psi_{\text{solid_effec}} - \psi_{\text{shadedleaf}}) \tag{2}
$$

$$
K_{stem} = Sapflow/(\psi_{shadedleaf} - \psi_{sunlitleaf})
$$
\n(3)

$$
K_{soil_plant} = Sapflow / (\psi_{soil_effec} - \psi_{sunilitleaf})
$$
\n(4)

260 During one measurement day, four values of the Ksoil_root, Kstem, and Ksoil_plant were obtained from measurements between 11AM and 2 PM. The average and standard deviation of these hourly measurements were calculated for each measurement day in order to present the seasonal dynamics of those variables. To capture the diurnal and seasonal variations of sap flow and sunlit leaf water potential, in addition,we plotted the hourly sap flow and hourly difference of effective soil water potential and sunlit leaf water potential for three measurement days starting from predawn and whole seasons, respectively, 266 to derive the slope which is also $K_{\text{soil_plant}}$.

267 **2.5. Statistical analysis**

268

12 269 Regression analysis was performed to understand the relationship between the sap flow volume and the 270 difference of effective soil water potential and sunlit leaf water potential as well as the relationship 271 between the total aboveground biomass and cumulated water transpired (sap flow volume). These 272 analyses allow to derive the slope as proxy of K_{soil_plant} and transpiration use efficiency, respectively. Since 273 all measured data have their own measurement errors, the generalized Deming regression was employed. We performed relationships (via correlation coefficient and statistical significant levels) of midday leaf An, 275 Gs, and E with midday K_{stem}, K_{soil_plant, K_{soil_root}, sunlit leaf potential, ψ_{soil_effec}, and the difference of ψ_{soil_effec}} and sunlit leaf water potential (ψdifference). All data processing and analysis were conducted using the R statistical software (R Core Team, 2022).

3. Results

3.1. Root growth under different water treatments, soil types and climatic conditions

280 Observed root length (cm cm⁻²) from the minirhizotubes in different soil depths at the first week of June (stem elongation), around silking, and at harvest in two growing seasons are shown in the Figure 2. Root length was similar among water treatments at the start of stem elongation in both years (Fig. 2a & 2d). 283 The difference in root length was pronounced at silking and harvest between the soil types. More root growth was observed in the silty soil compared to the stony soil with the same water treatment (i.e. 2.5 - 6 times higher at depth 40 cm). This indicated the strong negative effects of stone content on root development. In the stony soil, root length in the irrigated plot (F1P3) was slightly higher than in the rainfed plot (F1P2). In contrast, the rainfed treatment (F2P2) in the silty soil showed much higher root length, especially from 40 to 120 cm depths as compared to the irrigated plot (F2P3) in both growing seasons. Much lower stone content and deep soil cracks in the silty soil (Morandage et al., 2021) allow root extension to the subsoil, particularly in the rainfed plot F2P2. Root length in the rainfed treatment (F2P2) in 2018, is higher than in 2017 which implies that root further developed to exploit the water in the soil under the rainfed condition to meet the higher evaporative demand.

[Insert Figure 2 here]

 294 Total root length (m m⁻²) estimated from minirhizotubes and its ratio to shoot dry matter (m kg⁻¹) at three measured dates (as in Figure 2) are shown in the Figure 3. Total root length was much higher for the silty 296 plots as compared to stony plots. In 2017, the highest total root length was observed in the rainfed plot of 297 the silty soil (F2P2) with approximately 9166 m m⁻² and 9878 m m⁻² around silking and harvest, respectively,

 which was almost two times higher than in the irrigated plot (F2P3). These figures were higher in 2018 299 than 2017 where total root length of F2P2 was 10188 m m⁻² and 13750 m m⁻² at silking and harvest time, respectively. For the rainfed stony soil (F1P2), soil water depletion around the beginning of June in 2017 (Supplementary material 1a) and from the first two weeks of June to harvest in 2018 (Supplementary material 2a) caused the strong reduction of shoot biomass. In the stony soil, the shoot dry matter of the 303 irrigated plot (F1P3) and the rainfed plot (F1P2) were 1275 and 536 g m⁻² at silking time (e.g. 19 July 2018 – DOY 200, Supplementary material 3a and 3b). However, there was a minor difference between F1P2 and F1P3 in terms of the ratio of root length to shoot dry matter. In the silty soil, a decrease of soil water potential was not pronounced (compared to stony soil) in both years 2017 and 2018 (Supplementary material 1b and 2b). In 2018, shoot biomass in the irrigated stony soil (F1P3) and silt soil (F2P3) were 308 similar (1275 and 1299 g m⁻², respectively on 19 July 2018 – DOY 200) while the shoot biomass of the 309 rainfed silty soil (F2P2) was 876 g m⁻² (Supplementary material 3a & 3b). However, the ratios of root length to shoot biomass in the rainfed plot of the silty soil (F2P2) were 3 and 6 times higher than those in the irrigated silty soil (F2P3) and stony soil (F1P3), respectively (e.g. 18 July, DOY 199). Moreover, total root 312 length was relatively equal among treatments at the start of set elongation (8 June - DOY 159, first week 313 of June) in both years, while this was the opposite for the ratio of root length to shoot dry matter. This firstly illustrated that the finer soil texture without stones and with soil cracks could favor the root growth which indicates strong interactions of root and soil conditions. Secondly, the larger root length and higher atmospheric evaporative demand in 2018 than 2017 indicates also the interaction of root growth and climatic conditions.

[Insert Figure 3 here]

319 **3.2. Stomatal conductance, photosynthesis, transpiration, and K_{soil plant}**

 3.2.1. Diurnal course of stomatal conductance, photosynthesis, transpiration, and water potential at leaf level

343 Seasonal stomatal conductance (Gs) and leaf water potential (ψ_{leaf}) are described in Figure 75 . The relationship between two variables was rather noisy and non-linear. The leaf water potential showed

345 distinct patterns among treatments in one growing season. Minimum ψ_{leaf} was maintained at around -1.5 346 MPa in the irrigated plot in stony soil (F1P3) and two plots in the silty soil (F2P2 and F2P3). Lower minimum 347 ψ_{leaf} could be observed in the rainfed plot with stony soil (F1P2) but it did not go beyond -2 MPa. Minor 348 leaf curling was observed only in the second week of June in the F1P2 in 2017. In 2018, the higher 349 temperature and vapor pressure deficit resulted in lower minimum ψleaf in all treatments and soil types as 350 compared to 2017. The minimum ψ_{leaf} was around -2 MPa in F1P3, F2P2, and F2P3 while ψ_{leaf} could drop 351 below -2 MPa in F1P2 which was due to the severe soil water deficit. The low Gs and ψ_{leaf} associated with 352 measurement dates when the substantial leaf curling was observed at mid of July to the end of growing 353 season in F1P2 in 2018 (Supplementary material 3c & 3d and Supplementary material 46c & d).

354 [Insert Figure 7-5 here]

16 355 The effective soil water potential (ψ_{soil_effect MD}), sunlit leaf water potential (ψ_{sunlitleaf MD}), stomatal 356 conductance (GS_{MD}), and whole plant hydraulic conductance (K_{soil_plant MD}) at midday at several times during 357 the growing season are presented in Figures 8-6 and 9-7 for 2017 and 2018, respectively. As expected, 358 there was not much difference in terms of $\psi_{\text{solid eff}}$ among F1P3, F2P2, and F2P3 from 02 August to one 359 week before harvest in 2017. The lowest $\psi_{\text{solid effec MD}}$ was observed in the F1P2. Leaf water potential 360 dropped drastically but also K_{soil_plant MD} increased strongly whereas ψ_{soil_effec MD} remained quite similar (e.g. 361 18 July). This is because sap flow have increased substantially in this day (e.g. from 2.34 mm d⁻¹ on 17 July 362 to 6.97 mm $d⁻¹$ on 18 July for the F1P2). The stomatal conductance decreased a lot in this day which could 363 be explained that the atmospheric demand increased (e.g. global radiation was 13.6 MJ m⁻² on 17 July 364 compared to 23.9 MJ on 18 July while daily VPD was 0.7 kPa and 1.2 kPa, respectively) even more than the 365 sap flow. Midday sunlit leaf water potential was not distinctively different among treatments with the 366 lowest ψ_{sunlitleaf MD} around -1.6 MPa throughout season. Also, Gs_{MD} was rather similar among plots. The 367 Ksoil_plant MD ranged from 0.125 to 0.96 mm h^{-1} MPa⁻¹ with a sharp reduction before harvest. In general, the 368 lowest values of K_{soil_plant MD} were found in F1P2 which was consistent with the smaller overall seasonal 369 Ksoil_plant (as the slope of linear relationship between sap flow and difference of effective soil water potential 370 and sunlit leaf water potential) (see Supplementary material 57).

371 [Insert Figure 86 here]

372 The ψ_{soil_effec MD} was substantially different in the two soil types and water treatments in 2018 (Figure 9a_{7a}). 373 Both F1P2 and F1P3 showed a gradual drop of ψ_{soil_effec MD} from 15 June until the third week of July then 374 increased after irrigation events on 18 July (Supplementary material 2b). However, ψ_{soil effec MD} of F1P2 was 375 much lower than F1P3 toward the harvest. The ψ_{soil_effec MD} of F2P2 and F2P3 only decreased progressively 376 from around 10 July till harvest even though there was water supply from the irrigation (Supplementary 377 material 2b). The water applied by irrigation and coming in by rainfall were insufficient to wet up the 378 deeper soil layers which remained dry. The low Gs_{MD} was corresponding to the lowest $\psi_{\text{sunittleaf MD}}$ and 379 K_{soil_plant MD} from the F1P2 (Figure $96 - 7c$ & $9d7d$). The K_{soil_plant MD} from all plots was ranging from 0.12 to 0.91 380 mm h⁻¹ MPa⁻¹. There was the drop in K_{soil_plant MD} (i.e. 3 to 9 July or 17-18 July) before irrigation in this plot. 381 However, it increased after the irrigation (i.e. 10 July and 19 July). This suggests that Ksoil_plant depends 382 strongly on the soil water content and the conductivity of the rhizosphere.

383 [Insert Figure 9-7 here]

384 **3.2.3. Relationships of stomatal conductance, transpiration, photosynthesis with plant hydraulic** 385 **variables at the plant canopy level**

17 386 The slope of linear relationship between sap flow and difference of ψ_{solid} effec and $\psi_{\text{sunlittleaf}}$ is shown for three 387 consecutive days (leaf water potential measurements from the predawn) and before and after irrigation 388 applications (17, 18, and 19 July 2018 or DOY 198, 199 and 200, respectively) (Figure 810). On both DOYs 389 dates 198-17 and 18 Julyand 199, the difference between ψ_{soil_effec} and ψ_{sunlitleaf} was around -1.6 MPa with 390 very low transpiration rates in the treatment F1P2 which was associated with very low plant hydraulic 391 conductance and leaf curling. The whole plant hydraulic conductance was disrupted on these two days 392 (0.06 and 0.16 mm h⁻¹ MPa⁻¹ for 17 and 18 JulyDOY 198 and 199, respectively). Water was supplied on DOY 393 18 July 199 at 1 PM for the irrigated plots (F1P3, F2P3) as well as F1P2 at 4 PM (for saving plant from death 394 due to severe drought stress). K_{soil_plant} was slightly changed (0.43 and 0.57 mm h⁻¹ MPa⁻¹ for F1P3 on DOY 395 199-18 and 19and 200 July, respectively and 0.5 and 0.58 mm h⁻¹ MPa⁻¹ for F2P3 on 18 and 19 JulyDOY 199 396 and 200, respectively). However, the increase of K_{solid} _{plant} was substantial in the F1P2 after the irrigation. 397 Soil water replenishment and an increase in the root - soil contact (Fig. $9a\overline{7a}$) allowed the K_{soil_plant} to 398 recover overnight to 0.46 mm h^{-1} MPa⁻¹. This resulted in a narrower water potential gradient between root 399 zone and sunlit leaf and in a higher transpiration rate on 19 JulyDOY 200.

400 [Insert Figure 408 here]

401 Seasonal average of different midday hydraulic conductance components (root system hydraulic 402 conductance - $K_{\text{sol_root}}$, stem hydraulic conductance – K_{stem} , and whole plant hydraulic conductance – 403 K_{solid_plant} are shown in Figure 449 . In the same year, the K_{stem} was not much different among F1P3, F2P2, 404 and F2P3 plots. The K_{stem} of those plots was slightly higher than in the F1P2 in both years. In general, the 405 Ksoil_root was lower than the Kstem. Overall, the estimated Ksoil_plant was around $1/$ ($1/K_{\text{soli_root}} + 1/K_{\text{stem}}$) 406 regardless of soil types, years, and water treatments. The K_{soil_root} and K_{soil_plant} in the F1P2 in 2018 was much 407 lower than the remaining plots while the K_{soil_root} and K_{soil} _{plant} was were not much different among plots in 408 2017. Our results indicated that there was an impact of soil hydraulic conductance on K_{soil root} and K_{soil plant.} 409 The Ksoil_plant and Ksoil_root depend strongly on the soil water content and the soil hydraulic properties. Overall, 410 the estimated K_{soil plant} was around 1/ (1/K_{soil root} +1/K_{stem}) regardless of soil types, years, and water 411 **treatments.** Although there is a large difference in total root length between the two soil types (e.g. F1P3 412 versus F2P2 or F2P3 versus F2P2), K_{soil_root} and K_{soil_plant} in those two plots were not much different. This 413 could be explained by the fact that K_{soil plant} was not only depended on root length but also depended on 414 the variability of root segment hydraulic conductance. The K_{soil}_plant and K_{soil}_pour depend strongly on the soil

415 water content and the soil hydraulic properties. Therefore

416 property but also a soil property.

417 [Insert Figure 449 here]

3.3. Relative importance of root and leaf area growth to transpiration and crop performance at canopy level

 Drought stress was observed in the rainfed plot (F2P2) in the second week of June 2017 with mild leaf rolling. The crop then recovered due to sufficient rainfall and lower evaporative demand. Drought stress occurring again at the stem elongation phase caused reduction of plant size (height and stem diameter) (Supplementary material 46) as well as a slight reduction of leaf area and biomass in this plot (Supplementary material 3a & 3c). Transpiration per unit of leaf area did not differ much among water 425 treatments and soil types in 2017 (Figure Supplementary material S842). The opposite was the case for the transpiration rate per unit of root length. The observed root length at different soil depths (Figure 2) and total root length for two plots in the stony soil was much smaller than in the silty soil (Figure 3). Therefore, transpiration per unit of root length in the stony soils (F1P2 & F1P3) was almost 3 times higher than transpiration in the silty soil. For the same soil, transpiration per unit root length of the irrigated treatment was slightly larger than in the rainfed plot.

[Insert Figure 12 here]

 The differences in sap flow per plant between water treatments and soil types were more pronounced in 433 2018 (Figure Supplementary material S439). The highest transpiration rate was observed in the irrigated plots (F1P3 & F2P3), followed by the rainfed plot of the silty soil (F2P2) and it was lowest in the rainfed plot of the stony soil (F1P2). These observations were in line with the differences in biomass and leaf area index between the treatments (Supplementary material 3b & 3d) and plant size (Supplementary material $64b-c-d$). In 2018, severe leaf rolling was observed in the rainfed plot (F1P2) from the beginning of June

 until the end of the growing period in 2018 (Supplementary material 3d). Similar to 2017, transpiration 439 per unit of root length was much higher in the stony plots as compared to silty plots. Also, for the silty soil, transpiration per unit of root length of the irrigated plot (F2P3) was higher than in the rainfed plot (F2P2). **[Insert Figure 13 here]** Higher cumulative transpiration in the irrigated plots did not result in higher transpiration use efficiency

443 (TUE) in both soil types (Figure). For instance, TUE were 16.87 g mm⁻¹ and 15.59 g mm⁻¹ for F1P2 and 444 F2P2, respectively, while they were 15.47 and 14.79 g mm⁻¹ for F1P3 and F2P3, respectively, in 2017 (Figure 445 104aA). For the same soil, the rainfed plot showed slightly higher TUE than the irrigated plot. When comparing the TUE of maize of the two soil types for the same water treatment, TUE at the stony soil was almost the same in silty soil. The TUE was not much different among treatments and soil types in 2018. 448 Overall, TUE in 2017 was higher as compared to 2018 (Fig. 104b).

449 [Insert Figure 14-10 here]

4. Discussions

4.1. Effects of soil types, water application, and climatic condition on root growth

 Our root observationsshowed that soil type considerably affected root growth more than water treatment (Figure 2). Root growth was strongly inhibited by the stony soil where much lower root length was observed than in the silty soil, especially in the deeper soil layers. This was consistent with the findings reported in (Morandage et al., 2021) where a linear increase of stone content resulted in a linear decrease of rooting depth across all stone contents and developmental stages. Also, both simulations and 457 observations indicated that rooting depth was sensitive increased due to the presence of cracks in the lower minirhizontron facility (Morandage et al., 2021) which could explain the high root length between 459 40 and 120 cm soil depths which was observed in the silty soil in both years. In the silty soil, root growth 460 was favored towards deeper soil layers as also reported for the same field in 2016 for winter wheat

461 (Nguyen et al., 2020). Observation in field grown maize, the higher root length density and root diameter 462 were found in the sand than in the loam. This was attributed to the higher investment in nutrient 463 exploration because the lower concentration of plant-available nutrients was in sand than in loam 464 (Vetterlein et al., 2022). Also, the larger root diameters in sand than in loam are more likely explained by 465 the need for soil contact of the roots (Jorda et al., 2022; Vetterlein et al., 2022).

466 Our total root length was in the reported range of Cai et al., (2018) who studied winter wheat roots on the 467 same soil types in 2016. The total root length in our work was higher than the reported results from Cai 468 et al., (2018) especially in the rainfed plot of the silty soil (F2P2) in 2018 (Fig. 3). In terms of the root: shoot 469 ratio, our observations were in line with those reported in the same soil types for wheat in Cai et al., (2018). 470 Grdóñez et al., (2020) has reported much larger figures of for instance 880 cm g⁻¹ in different locations and 471 under different N application rates in maize growing in the Midwest of US. Jorda et al., (2022) reported a 472 wide range of ratios of root length to shoot biomass root: shoot ratio-from 200 to 1000 cm g⁻¹ around 473 flowering time of maize depending on the wild type and root hair mutant genotypes growing on either 474 loamy or sandy soils. More roots and higher ratios of root length to shoot biomassroot: shoot ratios were 475 found in the sand than in the loam in both wild type and root hair mutant genotypes (Jorda et al., 2022; 476 Vetterlein et al., 2022). Cai et al., (2018) observed much larger ratios of root length to shoot biomass root: 477 shoot ratio in drought stressed plots than in irrigated plot in both soil types in winter wheat which 478 indicated the alternation of sink: source relationships to cope with water stress. This study emphasized 479 that more assimilates are used to promote root growth and extract more water under drought stress. 480 However, this was not the case for the stony soil in our work where the drought stress was more 481 pronounced, especially in 2018. A slightly higher root: shoot ratio in the F1P2 treatment compared to F1P3 482 (DOY 194 & 255) was observed in 2017 while the root: shoot ratio in the two treatments was almost the 483 same on DOY 199 and 228 in 2018 (Fig. 3). We only observed much higher root: shoot ratio in the rainfed 484 plot (F2P2) as compared to the irrigated plot on the silty soil (F2P3). Comas et al., (2013) has reported that

 In our study, stomata closed earlier and at more negative soil and leaf water potentials in stony soil than 504 in silty soil (see Fig. 4, 5, 6 and 7). In other the previous work, Koehler et al., (2022) reported that maize stomata closed at lower negative leaf water potentials in sand than in loam growing under controlled environment. Cai et al., (2022b) investigated transpiration response of pot-grown maize in two contrasting soil textures (sand and loam) and exposed to two consecutive VPD levels (1.8 and 2.8 kPa). Transpiration rate decreased at less negative soil matric potential in sand than in loam at both VPD levels. In sand, high

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4.2.2. Hydraulic conductance components as affected by soil water conditions

 Estimates of hydraulic components in soil-plant-atmosphere continuum are important not only to 553 understand its underlying relationship to other crop characteristics (stomatal conductionance, transpiration, and photosynthesis) but also to provide modeling parameters in process-based soil-root- shoot models (Nguyen et al., 2020; Sulis et al., 2019; Nguyen et al., 2022b). Measurement of the components of hydraulic conductance are challenging under field conditions because it requires the

variables

 In 2017, our estimated midday effective soil water potential (ψsoil_effec MD) did not vary much (between soil types and treatments) which was consistent with the low variability in midday sunlit leaf water potential

615 The transpiration rate and K_{soil_plant} (slope of linear regression lines in Fig. $480a$ and b) were very low in the 616 rainfed plot under the stony soil (F1P2) which was associated with the large ψ_{difference} (Fig. 10a 8a & b) and 617 the lower stomatal conductance as compared to other plots (Fig. 97c). The K_{soil_plant} slightly increased after 618 irrigation (18 July - DOY 199 in Fig. 810b) corresponding with the smaller $\psi_{\text{difference}}$ (Fig. 180b) and an 619 increase in stomatal conductance (Fig. $79c$). Seasonal K_{soil_plant} was low in the rainfed plot under stony soil 620 (F1P2) with the larger $\psi_{difference}$ (Supplementary material 57). In addition, our study showed that the midday 621 stomatal conductance, photosynthesis, and transpiration were significantly correlated only with midday 622 Ksoil_plant in the rainfed plot on the stony soil (F1P2) in 2018 where high VPD and temperature occurred 623 (Supplementary material 610, 117, and Supplementary material 812). Maize plants had lower plant 624 hydraulic conductance and more negative soil water potential in the rainfed plot in stony soil required the 625 larger gradients in soil water pressure to sustain the same transpiration rate (thatthus and they exhibited 626 earlier stomatal closure) as compared to the same plot in the silty soil. This was in line with a study from 627 Abdalla et al., (2022) which suggested that during soil drying, stomatal regulation of tomato is controlled 628 by root and soil hydraulic conductance. Recent work from Müllers et al., (2022) on faba bean and maize suggested that differences in the stomatal sensitivity among plant species can be partly explained by the sensitivity of soil-plant hydraulic conductance to soil drying. The loss of conductance has immediate consequences for leaf water potential and the associated stomatal regulation. Cai et al., (2022b) also showed that the decrease in sunlit leaf stomatal conductance was well correlated with the drop in soil- plant hydraulic conductance, which was significantly affected by soil texture. This was confirmed in our 634 work where the stony soil strongly impacted on root growth, modulated $K_{\text{sol_plant}}$, and consequently influenced the leaf stomatal conductance, photosynthesis, and transpiration.

 4.3. Relative contribution of water control by leaves and roots on transpiration and transpiration use efficiency

 Responses of crops via stomatal control to reduce water loss at leaf scale while maintaining leaf photosynthesis and water use efficiency were reported earlier (Nguyen et al., 2022a; Vitale et al., 2007). 640 In addition to that, in the maize experiments in 2017 and 2018In our study, leaf rolling was observed in both rainfed plots on the stony and the silty soil in the second week of June 2017 and from the beginning of June until the end of the growing period in 2018. This indicates another dehydration avoidance mechanism resulting from morphological adjustments which is an effective mechanism for delaying senescence (Aparicio-Tejo and Boyer, 1983; Richards et al., 2002). Stomatal closure resulted in more reduction of transpiration and assimilation in the rainfed plots than irrigated plots with the same soil type 646 (Fig. 54, Supplementary material Fig. 64 & 5, Fig. 75, and Fig. 13Supplementary material 9Aa). There was reduction of shoot biomass (also stem size and leaf size adjustments) in F1P2 as compared to other plots. However, the TUE was not smaller in this plot than the remaining plots. These observations confirm that plant size adjustments through reduction of height, leaf width and length are efficient responsesto reduce water loss at canopy scale in addition to stomatal control at the leaf level.

 Relative contribution of leaf area to transpiration has been highlighted in wheat where reduction of tiller number resulted in significantly (lower LAI, thus lower canopy transpiration (Cai et al., 2018; Trillo and

678 in regulating plant transpiration (Cai et al., 2022a). Other traits like root hair density (Cai et al., 2022a) or 679 higher root length density (Vadez, 2014) could contribute to the soil to root water potential and root-zone 680 hydraulic conductance where dense root hairs are delayingdelayed soil water deficit in drying soils. 681 However, contrasting results have shown that root hairs did not have an effects on root water uptake (see 682 Jorda et al. 2022). The role of root hairs could not be analyzed in our work which was based on the root 683 data from minirhizotron images.

684 **5. Conclusion**

 We presented plant hydraulic characteristics and crop growth from root to shoot of maize under field- grown conditions with two soil types (silty and stony), each soil with two water regimes (irrigated and 687 rainfed) for two growing seasons (2017, 2018). Our results confirmed that root length and ratios of root 688 length to shoot biomassroot: shoot ratio was were modulated by soil types and water treatment but less 689 by seasonal evaporative demand. Increase ratio of root length to shoot biomassroot: shoot ratio has been an important response of maize that allows plants to extract more water under drought stress that occurred rather in the silty soil but less in the stony soil due to the higher content of stony material. Despite of lower root length in the stony irrigated plot, transpiration rate was not much lower than in the silty irrigated plot. This could be related to another property of the root such as root segment conductance or other root traits (e.g. root hair). Further investigation with extensive measurements of rootsincluding axial and radial root conductance at field scale will be required to better explain the observed results.

696 Another conclusion is that stomatal regulation maintains leaf water potential at certain thresholds which 697 depends on soil types, soil water availability, and seasonal atmospheric demand. The stomata conductance 698 was smaller and decreased at more negative leaf water potentials in stony soil than in silty soil. The leaf 699 water potentials are affected by the soil-plant plant hydraulic conductance. In addition to stomatal 700 regulation, leaf growth and plant size adjustments are important to regulate the transpiration that and 701 water use efficiency was not different among treatments and soil types in the same year.

 The lowest soil-plant hydraulic conductance was observed in the stony soil with severe drought stress as compared to silty soil while its variation depends also on the soil water variation (before and after irrigation). Root system and soil-plant hydraulic conductance depended strongly on soil hydraulic properties. In the stony soil, which has a considerably smaller water holding capacity than the silty soil, root length was considerably smaller than in the silty soil. Nevertheless water uptake per unit root length was much larger than in the fine soil. This also means that the hydraulic conductance per unit root length must have been much larger in the stony soil than in the fine soil. Cai et al., (2018) observed a similar effect for winter wheat but they found much smaller differences in the root length normalized root conductance. The higher root length normalized root conductance means that the anatomy of the root tissues must have been influenced by the soil texture and compensated the considerably smaller root length in the stony soil. Looking at the effect of water treatments in the silt soil, the non-irrigated plot had more roots than the irrigated one and both had more roots in the year with high VPD. But the soil-root conductance was higher in the irrigated plot than in the rainfed plot. This means that in the irrigated plot, the soil-root conductance per unit root length was higher than in the rainfed plot. This could either be due to wetter soil conditions and higher soil conductance or it could be due to a larger conductance of the root tissues. Especially in 2017 when the silty soil was wetter, the slightly larger soil-root conductance in the irrigated plot is most likely the result of larger root tissue conductance in the irrigated plot. Thus, how root architecture (here represented simply by the total root length) and root tissue conductivities 'respond' to drought stress might be opposite depending on the comparisons that are made. When the stony soil and silt soil are compared, the higher 'stress' due to lower water availability in the stony soil resulted in less roots with a higher root tissue conductance in the soil with more stress. When comparing the rainfed with the irrigated plot in the silty soil, the higher stress in the rainfed soil resulted in more roots with a lower root tissue conductance in the treatment with more stress. This illustrates that the 'response' to stress can be completely opposite depending on conditions or treatments that lead to the differences in stress that are compared. Therefore, it cannot be the 'stress' alone that defines how a plant will react and adapt its

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758 **List of Tables**

759 Table 1. Crop phenology and management information for different treatments in 2017 and 2018.

Notes: ¥ 760 Notes: $*$ from sowing to harvest; $*$ for rainfall for whole growing season;

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Author contribution

 Huu Thuy Nguyen, Thomas Gaiser, Jan Vanderborght, and Frank Ewert: Conceptualization; Huu Thuy Nguyen, and Hubert Hüging: Data curation and data quality check (aboveground measurements); Lena Lärm, Felix Bauer, Anja Klotzsche, Jan Vanderborght, and Andrea Schnepf: data curation and data quality check (belowground measurements); Huu Thuy Nguyen: Formal data analysis and visualization; Thomas Gaiser, Jan Vanderborght, Andrea Schnepf, and Frank Ewert: Funding acquisition & Project administration; Huu Thuy Nguyen: writing – original draft; all authors: review, editing, and finalizing the manuscript.

Competing interests

 This manuscript has not been published and is not under consideration for publication in any other journal. All authors agreed and approved the manuscript and its submission to this journal. We declare there is no conflict of interest.

Code/Data availability

List of Figures

Figure 1: Daily maximum air temperature (Tmax) (°C), daily maximum air vapor pressure deficit (VPD) (kPa) in the two growing seasons (a) 2017 and (b) 2018 and cumulative (sum) of rainfall and irrigation from the rainfed (P2) and irrigated (P3) plots of the stony soil (F1) and silty soil (F2) in the two growing seasons (c) 2017 and (d) 2018. The black dashed vertical lines (a) and (b) indicate silking time. Grey vertical lines in (a) and (b) indicate the measured days for leaf gas exchange and leaf water potential. Two lines for 2017F2P2 and 2017F2P3 were overlapped by the lines from 2017F1P2 and 2017F1P3, respectively

Figure 2: Observed root length from minirhizotubes (cm cm⁻²) from 10, 20, 40, 60, 80, and 120 cm soil depth from the rainfed (P2) and irrigated (P3) plots of the stony soil (F1) and silty soil (F2) in the two growing seasons in 2017 (a - 8 June, b - at silking on 13 July, c - at harvest on 12 September) and in 2018 (d - 7 June, e - at one week after silking - 18 July, f - one week before harvest - 16 August).

Figure 3: Observed root length from minirhizotubes (m m^{-2}) and ratio of root length per shoot dry matter (m $kg⁻¹$) from the rainfed (P2) and irrigated (P3) plots of the stony soil (F1) and silty soil (F2) in the two growing seasons (DOY 159, 194, and 255, left panel) in 2017 and in 2018 (DOY 158, 199, and 228, right panel) where on 8 June (DOY 159) at silking on 13 July (DOY194) 2017; and at harvest on 12 September (DOY 255) in 2017; 7 June (DOY 158), one week after silking on 18 July (DOY 199); and one week before harvest on 16 August (DOY 228) in 2018 (see also Figure 2).

Figure 4. Diurnal course of (a) photosynthetically active radiation (PAR) and vapor pressure deficit (VPD), (b –e) leaf net photosynthesis (An), (f –i) leaf stomatal conductance (Gs), (j –m) leaf transpiration (E), and (n –q) leaf water potential (LWP) on 18 July in maize in 2018 before irrigation at the rainfed (P2) and irrigated (P3) plots of the stony soil (F1) and silty soil (F2). Measurement was carried out from shaded leaf (plus symbol with line) and two sunlit leaves (solid dot - lines and solid square - lines). Crop was irrigated at 1 PM, 1 PM, 4 PM for F1P3, F2P3, and F1P2, respectively (22.75 mm for each plot) (Supp. 2). Black arrows indicate time of irrigation.

Leaf position $^{+}$ shaded sunlit

Figure 5: Seasonal stomatal conductance to water vapor (Gs) versus leaf water potential (ψ_{leaf}) in 2017 (top panel) and in 2018 (bottom panel) at the rainfed (P2) and irrigated (P3) plots of the stony soil (F1) and silty soil (F2). Vertically continuous and dashed lines indicated ψ_{leaf} at -1.5 and -2 MPa, respectively. Measurement was carried out from shaded leaf (plus symbol) and two sunlit leaves (solid dots)

Figure 6: Dynamic of around midday (MD) of (a) the effective soil water potential (ψ_{soil_effec, MD}) (b) sunlit leaf water potential (ψ_{sunlitleaf MD}), (c) stomatal conductance (Gs MD) and (d) whole soil-plant hydraulic conductance (K_{soil plant MD}) in the growing season 2017 from the rainfed (P2) and irrigated (P3) plots of the stony soil (F1) and silty soil (F2). Error bars indicate the standard deviation of the different values taken around midday (11 AM, 12AM, 1PM, and 2 PM) of different sunlit leaves. Whole soil-plant hydraulic conductance was shown from 17 July when sap flow was measured. The black arrows indicates the irrigation events for the irrigated treatments F1P3 and F2P3 in the showing period.

Figure 7: Dynamic of around midday (MD) of (a) the effective soil water potential (ψ_{soil_effec MD}) (b) sunlit leaf water potential (ψ_{sunlitleaf MD}), (c) stomatal conductance (Gs MD) and (d) whole soil-plant hydraulic conductance (K_{soil_plant MD}) in the growing season 2018 from the rainfed (P2) and irrigated (P3) plots of the stony soil (F1) and silty soil (F2). Error bars indicate the standard deviation of the different values taken around midday (11 AM, 12AM, 1PM, and 2 PM) Leaf water potential and stomatal conductance were 2 sunlit leaves and one shaded leaf at each measured hour. Whole soil-plant hydraulic conductance was shown from 3 July when sap flow was measured. The black arrows indicates the irrigation events for the irrigated treatments F1P3 and F2P3 while the orange arrow indicates the irrigation application for the rainfed plot at the stony soil (F1P2).

Figure 8: Relationship of sap flow and difference of effective soil water potential and sunlit leaf water potential (ψ_{difference}) from the rainfed (P2) and irrigated (P3) plots of the stony soil (F1) and silty soil (F2) on three consecutive measurement days from predawn in 2018 (a) 17 July - DOY 198, (b) 18 July - DOY 199 and (c) 19 July - DOY 200. Crop was irrigated on 18 July (DOY 199) at 1 PM, 1 PM, and 4 PM for F1P3, F2P3, and F1P2, respectively (22.75 mm for each plot). The unit of slope in the linear regression (or soil-plant hydraulic conductance) is mm h^{-1} MPa⁻¹. Regression was based on the DEMING approach. The asterisk which are next to the slopes indicate a significant correlation between two variables according to Pearson method (ns: non-significant; * p < 0.05; ** p < 0.01; *** p < 0.001).

Figure 9: Comparison of different midday hydraulic components (mm h⁻¹ MPa⁻¹): soil-plant (grey bars), soilroot (yellow bars), and stem (blue bars) from the rainfed (P2) and irrigated (P3) plots of the stony soil (F1) and silty soil (F2) in the two growing seasons (a) in 2017 and (b) in 2018. The error bars indicate the standard deviation from measurements around midday (11 AM, 12AM, 1PM, and 2 PM) in different measured days (in 2017 with n = 4 x 9 days, Supplementary material 10, 11, and Fig. 6 and in 2018 with n = 4 x 10 days, Supplementary material 10, 12, and Fig. 7).

Figure 10: Relationship of aboveground dry matter and cumulative sap flow from the rainfed (P2) and irrigated (P3) plots of the stony soil (F1) and silty soil (F2) in the two growing seasons (a) 2017 and (b) 2018. The unit of slope linear relationship is g mm⁻¹. The less number of data points in (b) in 2018 from the F2P2 and F2P3 plots were due to the missing values of measured sap flow because of sensor disconnection. For aboveground dry matter, each point represents the average of two sampling replicates, except the harvest with 5 sampling replicates.